FILE 'HOME' ENTERED AT 12:54:08 ON 23 FEB 2004

=> fil reg FILE 'REGISTRY' ENTERED AT 12:54:16 ON 23 FEB 2004 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2004 American Chemical Society (ACS)

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 22 FEB 2004 HIGHEST RN 652965-05-4 DICTIONARY FILE UPDATES: 22 FEB 2004 HIGHEST RN 652965-05-4

TSCA INFORMATION NOW CURRENT THROUGH JULY 14, 2003

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

*** YOU HAVE NEW MAIL ***

'.REGISTRY' IS DEFAULT FORMAT FOR 'REGISTRY' FILE

=> str
:=> d sia
L1 HAS NO ANSWERS
L1 STR

C~~C @22 @23

REP G1=(0-10) 22-12 23-14 VAR G2=N/S/O NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:
RSPEC I
NUMBER OF NODES IS 2

STEREO ATTRIBUTES: NONE

=> s 11

SAMPLE SEARCH INITIATED 13:03:09 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 235 TO ITERATE

100.0% PROCESSED 235 ITERATIONS

5 ANSWERS

108 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS: 3781 TO 5619
PROJECTED ANSWERS: 5 TO 234

L2 5 SEA SSS SAM L1

=> d scan

L2 5 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN

IN 2,4-Undecadienamide, 11-(1,3-benzodioxol-5-yl)-N-(cyclohexylmethyl)-, (E,E)- (9CI)

MF C25 H35 N O3

Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> s l1 ful

L3

FULL SEARCH INITIATED 13:03:54 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 3982 TO ITERATE

100.0% PROCESSED 3982 ITERATIONS

SEARCH TIME: 00.00.01

108 SEA SSS FUL L1

=> d tot reg 1 RN442159-02-6 REGISTRY 2 RN351432-51-4 REGISTRY 3 RN 351432-47-8 REGISTRY 4 RN290328-98-2 REGISTRY 5 RN228873-78-7 REGISTRY 6 RN219324-57-9 REGISTRY 7 RN219324-56-8 REGISTRY 8 RN 190592-66-6 REGISTRY 9 RN190592-65-5 REGISTRY 10 190592-64-4 REGISTRY RN 11 190592-63-3 REGISTRY RN12 RN190592-57-5 REGISTRY 13 190592-56-4 REGISTRY RN

C~~C @22 @23

REP G1=(0-10) 22-12 23-14 VAR G2=N/S/O :end

L4 STRUCTURE CREATED

=> search 14

ENTER TYPE OF SEARCH (SSS), CSS, FAMILY, OR EXACT:.
ENTER SCOPE OF SEARCH (SAMPLE), FULL, RANGE, OR SUBSET: sub
ENTER SUBSET L# OR (END):13
ENTER SUBSET SEARCH SCOPE - SAMPLE, FULL, RANGE, OR (END): ful
FULL SUBSET SEARCH INITIATED 13:06:42 FILE 'REGISTRY'
FULL SUBSET SCREEN SEARCH COMPLETED - 108 TO ITERATE

100.0% PROCESSED 108 ITERATIONS SEARCH TIME: 00.00.01

1 ANSWERS

L5 1 SEA SUB=L3 SSS FUL L4

=> d sub bib abs

L5 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2004 ACS on STN

RN 442159-02-6 REGISTRY

CN 2,4,13-Pentadecatrienamide, 15-(1,3-benzodioxol-5-yl)-N-(2-methylpropyl)-, (2E,4E,13E)- (9CI) (CA INDEX NAME)

OTHER NAMES:

CN Brachystamide C

FS STEREOSEARCH

MF C26 H37 N O3

SR CA

LC STN Files: ANABSTR, CA, CAPLUS

Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1907 TO DATE)
1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1

AN 137:90938 CA

TI Amides from Piper brachystachyum and Piper retrofractum

AU Banerji, Avijit; Sarkar, Manjusha; Datta, Ratna; Sengupta, Piyali; Abraham, Koshy

CS Centre of Advanced Studies on Natural Products, Department of Chemistry, University College of Science, Calcutta, 700 009, India

SO Phytochemistry (2002), 59(8), 897-901 CODEN: PYTCAS; ISSN: 0031-9422

B Elsevier Science Ltd.

PB Elsevier Scier DT Journal

LA English

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QX 861. P45

Three unsatd. amides, designated brachystamides-C (e.g. I), D and E have been characterized from Piper brachystachyum Wall. Brachystamide-C, shown to be N-isobutyl-15-(3',4'-methylenedioxyphenyl)-2E,4E,13E-pentadecatrienamide, was unusual in having a non-conjugated double bond. Piper retrofractum Vahl. yielded retrofractamide-D, which has been fully characterized.

Ι

RE.CNT 8 THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> fil beil

FILE 'BEILSTEIN' ENTERED AT 13:08:30 ON 23 FEB 2004 COPYRIGHT (c) 2004 Beilstein-Institut zur Foerderung der Chemischen Wissenschaften licensed to Beilstein GmbH and MDL Information Systems GmbH

FILE RELOADED ON OCTOBER 20, 2002 FILE LAST UPDATED ON DECEMBER 15, 2003

FILE COVERS 1771 TO 2003.

*** FILE CONTAINS 8,861,754 SUBSTANCES ***

>>> PLEASE NOTE: Reaction data and substance data are stored in separate documents and can not be searched together in one query.

Reaction data for BEILSTEIN compounds may be displayed immediately with the display codes PRE (preparations) and REA (reactions). A substance answer set retrieved after the search for a chemical name, a molecular formula or a structure search for example can be restricted to compounds with available reaction information by concatenation with PRE/FA, REA/FA or more general with RX/FA. The BEILSTEIN Registry Number (BRN) is the link between a BEILSTEIN compound and belonging reactions. For more detailed reaction searches BRNs can be selected from substance answer sets and searched in the next step as reaction partner BRNs - Reactant (RX.RBRN) or Product BRN (RX.PBRN). After a search for reaction details substance documents associated with reactants or products may be retrieved by